Mutations in Multiple Domains Activate Paramyxovirus F Protein-Induced Fusion

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SER virus, a paramyxovirus that is closely related to simian virus 5 (SV5), is unusual in that it fails to induce syncytium formation. The SER virus F protein has an unusually long cytoplasmic tail (CT), and it was previously observed that truncations or specific mutations of this domain result in enhanced syncytium formation. In addition to the long CT, the SER F protein has nine amino acid differences from the F protein of SV5. We previously observed only a partial suppression of fusion in a chimeric SV5 F protein with a CT derived from SER virus, indicating that these other amino acid differences between the SER and SV5 F proteins also play a role in regulating the fusion phenotype. To examine the effects of individual amino acid differences, we mutated the nine SER residues individually to the respective residues of the SV5 F protein. We found that most of the mutants were expressed well and were transported to the cell surface at levels comparable to that of the wild-type SER F protein. Many of the mutants showed enhanced lipid mixing, calcein transfer, and syncytium formation even in the presence of the long SER F protein CT. Some mutants, such as the I310 M, T438S, M489I, T516V, and N529K mutants, also showed fusion at lower temperatures of 32, 25, and 18°C. The residue Asn529 plays a critical role in the suppression of fusion activity, as the mutation of this residue to lysine caused a marked enhancement of fusion. The effect of the N529K mutation on the enhancement of fusion by a previously described mutant, L539,548A, as well as by chimeric SV5/SER F proteins was also dramatic. These results indicate that activation to a fusogenic conformation is dependent on the interplay of residues in the ectodomain, the transmembrane domain, and the CT domain of paramyxovirus F proteins.

Fusion of the paramyxovirus envelope and the host cell membrane is brought about by the interaction of two surface glycoproteins, the hemagglutinin-neuraminidase protein (HN) and the fusion (F) protein. The F protein is synthesized as an inactive precursor, F0, that is posttranslationally cleaved by a host protease into two disulfide-linked subunits called F1 and F2. The cleavage of F is essential for virus-cell and cell-cell membrane fusion and also for virus infectivity (12, 21, 28). A well-conserved hydrophobic domain (fusion peptide) at the amino terminus of F1 is exposed by the cleavage (13, 18) and is considered to be involved in the fusion event (8, 22). Three heptad repeat (HR) domains are found in the F1 ectodomain (4, 9). HR1 is immediately adjacent to the carboxyl terminus of the fusion peptide, while HR2 is close to the transmembrane (TM) domain; HR3 is located between the HR1 and HR2 domains (9). The importance of these domains for fusion was shown by mutational analysis (26, 31) and also by the use of peptide analogues which inhibit fusion, presumably because they interfere with the conformational changes in the molecule that are necessary for fusion (10). Crystallographic analyses have shown that polypeptide fragments representing the HR1 domain of the F protein of simian virus 5 (SV5), Newcastle disease virus, or human respiratory syncytial virus form a trimeric coiled-coil structure to which three antiparallel helices of the polypeptide fragments representing the HR2 domain can bind (3, 5, 39). Similar six-helix bundle structures are also found in other viral fusion proteins and represent the final postfusion form of the protein (7, 14).

The cytoplasmic tail (CT) domain of some viral fusion proteins has also been shown to play a regulatory role in membrane fusion. Among the paramyxoviruses, F-protein CT truncations in Newcastle disease virus show highly reduced syncytium formation (30). Truncations in the CTs of the parainfluenza virus type 3 and SV5 F proteins abolished fusion activity (2, 37), whereas truncation of the CT of the human parainfluenza virus type 2 F protein did not affect its fusion activity (37). In SV5, deletion of the CT of the fusion protein inhibited fusion pore enlargement (6). The region of the C-terminal 16 amino acids of the envelope protein of murine leukemia virus, called the R peptide, is also known to be inhibitory to membrane fusion (11, 17, 29, 36). SER virus is a recently identified paramyxovirus, which is very closely related to SV5 but replicates without syncytium formation (34). Comparison of the CT sequences between the two viruses revealed the presence of a stop codon at amino acid 530 in the SV5 F CT, whereas an additional 22-amino-acid-extended CT is found in the SER F protein. Truncations or mutations in the CT domain of the SER virus F protein were found to enhance syncytium formation, indicating that the elongated CT interferes with membrane fusion in a sequence-dependent manner (33, 34). Besides the long CT domain, SER virus F differs from SV5 F by nine residues, six of which are in the ectodomain, one of which is in the TM domain, and two of which are in the CT domain.

Previous studies have demonstrated that SV5 strains differ in their requirements for corexpression of the HN protein to induce membrane fusion and that these differences can be
mapped to specific amino acid residues in the external domain of the F protein (15, 25). The WR strain F protein requires the coexpression of HN for fusion activity, whereas the W3a strain F protein exhibits HN-independent fusion; however, even for the W3a F protein, HN coexpression enhances F-mediated fusion (27). A proline residue at position 22 was demonstrated to be important for conferring HN-independent fusion as well as faster kinetics and a lower temperature for the fusion activity of the W3a F protein, and conversion of a serine at position 443 to proline further enhanced its fusion activity in the presence or absence of HN (25). The SER virus F protein contains proline residues at positions 22 and 443 in the external domain, thus corresponding to a highly fusogenic S443M mutant of SV5 described by Paterson et al. (25). However, we observed a complete lack of fusion by SER virus F but only a partial suppression of fusion activity in a chimeric SV5 F carrying the SER virus F CT (33). These results indicate a role of other amino acid differences between the SER virus and SV5 F proteins in conferring a more active fusion phenotype.

For the present study, we investigated the effects of the nine amino acids which differ between the SER virus and SV5 F proteins on lipid mixing, content mixing, and syncytium formation. We also studied the effect of these amino acid differences in the context of previously described mutants (33) which were found to enhance F-mediated membrane fusion.

MATERIALS AND METHODS

Cells, viruses, and vectors. HeLa T4 cells were maintained in Dulbecco’s modified minimal essential medium (DMEM) supplemented with 10% fetal calf serum (HyClone Laboratories, Logan, Utah). The recombinant vaccinia virus vTF7-3 and the wild-type (wt) vaccinia virus strain IHD-J were kindly provided by Bernard Moss (National Institutes of Health, Bethesda, Md.). The vaccinia virus stocks were propagated and titrated on CV-1 cells. Plasmid pGINT7 β-Gal was provided by Edward Berger (National Institutes of Health). The previously characterized pGEM-3-SER F and pGEM-3-SER HN plasmids (34) were expressed in HeLa T4 cells. The pGEM-3-SV5 F and pGEM-3-SV5 HN plasmids were described previously (34). A rabbit anti-SV5 antibody was a kind gift from R. A. Lamb (Northwestern University, Chicago, Ill.). SER virus was propagated in MDBK cells and virus titers were determined by a hemagglutination assay using guinea pig or chicken erythrocytes.

Generation of mutant F proteins. Mutations in the SER virus F gene were generated by use of a Quick Change site-directed mutagenesis kit (Stratagene, La Jolla, Calif.). The pGEM-SER F plasmid was used as the template and the two synthetic oligonucleotide primers designed to carry the mutation at the desired mutations and the absence of additional mutations. All constructs were sequenced to confirm the presence of the amino acid differences between the SER virus and SV5 F proteins.

Transfection, radiolabeling, and immunoprecipitation. The mutant and wt SER virus F proteins were expressed by using the vaccinia virus-bacteriophage T7 RNA polymerase transient expression system. Briefly, 35-mm-diameter dishes of subconfluent cells were infected with the vTF7-3 virus at a multiplicity of infection (MOI) of 10 for 1 h and were later transfected with 3 μg of plasmid DNA by the use of Lipofectin (Invitrogen, Carlsbad, Calif.). At 18 h posttransfection, the transfected cells were starved in DMEM lacking methionine and cysteine for 45 min, pulse labeled with 100 μCi of [35S]methionine-cysteine/ml for 30 min at 37°C, and then chased with DMEM containing 10% fetal calf serum for 2 h. The cells were washed thrice with ice-cold phosphate-buffered saline (PBS) containing 0.1 mM CaCl₂ and 1 mM MgCl₂ (PBS-CM) and then incubated with 1 ml of a 0.5-mg/ml solution of sulfo-L-cysteine (2-biotin-amido)ethyl-1,3-dithiopropionate (NHS-SS-biotin; Pierce) in PBS-CM at 4°C for 30 min. Free biotin was removed by a brief incubation with DMEM containing 10% fetal calf serum, and the cells were washed thrice with PBS-CM. The biotinylated cell surface proteins were lysed and immunoprecipitated with an anti-SV5 antibody (which also recognizes SER virus-encoded proteins) and a protein A-agarose beads (Pierce). The cell lysate-carrying beads were washed thrice and divided into two aliquots. One aliquot was used for immunoprecipitation and the other aliquot was boiled in 10 μl of 10% SDS and diluted with 1 ml of lysis buffer. The supernatant from the protein A-agarose beads was incubated with streptavidin-agarose beads for 2 h at 4°C. The proteins characterized by SDS–8% polyacrylamide gel electrophoresis and autoradiography. Quantitation of the surface expression of the S1 protein was performed by fluorescence-activated cell sorting (FACS) analysis. The results are presented as percentages of the cell surface expression levels obtained for the corresponding F proteins compared to that of the wt SER virus or SV5 F protein.

Pulse-chase analysis. Dishes (35-mm diameter) of subconfluent HeLa T4 cells were infected for 1 h with the vTF7-3 virus at an MOI of 10 and then transfected with wt and mutant SER virus F plasmids. At 18 h posttransfection, the transfected cells were starved in DMEM lacking methionine and cysteine for 45 min, pulse labeled with 100 μCi of [35S]methionine-cysteine/ml for 30 min at 37°C, and then chased with DMEM containing 10% fetal calf serum for 0, 15, 30, 45, 60, or 120 min. The cells were washed thrice with ice-cold PBS-CM at the indicated time points and incubated with 1 ml of a 0.5-mg/ml solution of NHS-SS-biotin (Pierce) in PBS-CM at 4°C for 30 min. The cells were washed and then lysed with cell dissociation buffer (10 mM Tris-HCl [pH 8.0], 250 mM NaCl, 0.5% Triton X-100, and 0.5% sodium deoxycholate) at the indicated time points. The surface expression of the SER virus wt and mutant F proteins was monitored by surface biotinylation studies as described above.

FIG. 1. Schematic diagram of SER virus and SV5 F proteins and mutants. (A) Differences in CT domains and presence of a stop codon at amino acid position 530 in the SV5 F CT followed by a 21-amino-acid coding sequence identical to that of the extended CT of the SER virus F protein. (B) SER virus F protein and nine amino acid differences between the SER virus and SV5 F proteins. The amino acid preceding the residue number denotes the wt SER virus F residue and the amino acid after the residue number denotes the SV5 F residue. L511A, a mutation of a conserved residue which served as a negative control, and mutants L539A and L548A, which were described earlier, are also shown (indicated with asterisks).
and suspended in PBS. HeLa T4 cells (10^5) were then analyzed with an antibody (1:100) followed by a FITC-conjugated secondary antibody, 5 mM EDTA in PBS, washed in PBS, incubated with an anti-SV5 mutant SER virus F protein or vTF7-3-expressing T7 polymerase as a T4 cells were infected with vTF7-3 at an MOI of 10 for 1 h at 37°C (A) Pulse-chase analysis of wt and mutant SER virus F proteins. HeLa T4 cells were infected with a vTF7-3 (MOI = 10) expressing wt or mutant SER virus F protein, and then transfected with neuraminidase before the two cell types were mixed. Cytosine arabinoside (40 μg/ml) was added to each plate. A second population of HeLa T4 cells was infected with wt vaccinia virus strain IHD-J and transfected with the pGINT7 β-Gal plasmid, which contains the β-galactosidase (β-Gal) gene under the control of the T7 promoter. At 16 h posttransfection, the two cell populations were collected and mixed by addition of 100 μl of each cell population to a flat-bottomed 96-well tissue culture plate. The plate was incubated at 37°C for 5 h, and cell fusion was measured by a colorimetric lysate assay as described previously. The quantity of β-galactosidase was calculated by comparing the hydrolysis rate for each sample with that of a standard commercial preparation of *Escherichia coli* β-galactosidase (Roche). The data were analyzed with the Delta Soft II Microplate analysis program. The fusion activity of the mutants was determined as a percentage of the β-galactosidase production observed in cells coexpressing wt SV5 F and HN.

**Temperature-dependent lipid mixing and calcine transfer assay.** Guinea pig erythrocytes (RBCs) were labeled with the hydrophobic fluorescent dye R18 (Molecular Probes, Eugene, Oreg.) as described by Bagai and Lamb (1). In brief, 15 μl of a 1 mg/ml solution of R18 in ethanol was added to 2 ml of 50% hematocrit fresh guinea pig RBC suspension in 10 ml of PBS that was being mixed on a vortex machine. After incubation at room temperature in the dark for 30 min, 35 ml of DMEM with 10% fetal calf serum was added to the suspension and further incubated for 20 min to adsorb the unbound probe. After being labeled, the RBC suspension was washed five times with 50 ml of PBS each time to remove the unincorporated R18. The last washing step was done with PBS containing an additional 1 mM CaCl_2 and 1 mM MgCl_2 (PBS+/−). The cells were resuspended to 50% hematocrit. For the preparation of doubly labeled RBCs, R18-labeled RBCs were loaded with calcine-AM (Molecular Probes, Leiden, Netherlands) as described by Melikyan et al. (20). R18-labeled RBCs (0.3 ml) were washed in swelling buffer (7.125 ml of PBS, 0.375 ml of H_2O). Calcine-AM was added to a 50% hematocrit RBC suspension to a final concentration of 5 μM, and the cells were washed and resuspended in 15 ml of PBS+/− (1% hematocrit). For determinations of lipid mixing and calcine transfer, BHK21 or HeLa T4 cell monolayers expressing SER virus F and HN or SER virus mutant F and HN and incubated with 50 μl of neuraminidase/ml in DMEM for 16 to 18 h posttransfection at 31°C were washed twice with PBS and first incubated with 1 ml of R18- and calcine-labeled RBCs (0.3% hematocrit) in PBS+/− at 4°C for 1 h in the dark with intermittent gentle agitation and then incubated at various temperatures for 15 min. Fusion was monitored and photographed by use of an inverted fluorescence microscope. Vaccinia virus-infected and mock-transfected cells were used as negative controls.

**RESULTS**

**Construction and expression of mutant proteins.** It was previously observed that the elongated cytoplasmic domain of the SER virus F protein is inhibitory to the process of membrane fusion (34). Serial truncations in the CT indicated the importance of an eight-amino-acid stretch between amino acids 535 and 542 in suppressing membrane fusion activity. Alanine scanning mutagenesis studies confirmed the importance of this amino acid sequence as well as a leucine residue at position 548 in the regulation of SER virus F-induced membrane fusion (33). We also recently obtained evidence that the presence of the long CT in the SER virus F protein stabilizes the metastable prefusion conformation of the F protein so that the conversion to the fusogenic conformation does not occur efficiently at a neutral pH or physiological temperature but may be triggered by exposure to a reduced pH or increased temperature. We observed that the lipid mixing, cytoplasmic content mixing, and syncytium formation ability of the SER virus F protein coexpressed with the SER virus HN protein were enhanced, both qualitatively and quantitatively, at elevated temperatures or under reduced pH conditions ranging between 4.8 and 6.2 (32). We also found that the fusion inhibition effect of the long CT of the SER virus F protein could be

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**Cell fusion assays.** Dishes (35-mm diameter) of subconfluent HeLa T4 cells were infected for 1 h with the vTF7-3 virus at an MOI of 10 and then transfected with 3 μg of a wt or mutant SER virus F construct with or without cotransfection of the wt SER HN plasmid by the use of Lipofectin. At 16 to 20 h posttransfection, the cells were monitored for syncytium formation under a light microscope.
A comparison of the nucleotide sequences of the F genes of SER virus (EMBL nucleotide sequence database; accession no. AJ278916) and the W3 strain of SV5 (24) revealed the presence of nine amino acid substitutions spanning the SER F protein. The nucleotide sequence differences pointed to nine amino acid substitutions that varied between SER virus and SV5 F proteins. Comparison of the nucleotide sequences of the F genes of SV5 and SER virus (EMBL nucleotide sequence database; accession no. AJ278916) and the W3 strain of SV5 (24) revealed the presence of nine amino acid substitutions spanning the SER F protein. These nucleotide sequence differences point to nine amino acid substitutions that varied between SER virus and SV5 F proteins.
virus F protein sequence (as shown in Fig. 1A) in addition to the presence of the long SER virus F CT. To determine the effects of the six ectodomains, one TM domain, and two CT differences on the lack of fusion by the SER virus F protein, we introduced these changes individually into the SER virus F sequence. The transport and stability of the mutant SER virus F proteins were examined by performing pulse-chase experiments. All of the mutants were found to be expressed at the cell surface and to have stability comparable to that of the wt SER virus F protein (Fig. 2A). The cell surface expression levels of the mutant proteins were estimated by FACS analysis, as shown in Fig. 2B. Except for the M370V mutant, which showed a surface expression level of 72% that of the wt SER virus F protein, all of the other SER virus F protein CT substitution mutants had surface expression levels of 88 to 96% that of wt F.

**Effect of mutations on lipid mixing and content mixing.** To investigate the effects of these mutations on different stages of membrane fusion, we transiently expressed the SER virus F and HN proteins in HeLa T4 cells in the presence of neuraminidase to prevent cell-cell fusion, and at 16 h posttransfection, we incubated the cells with RBCs that were doubly labeled with R18 and calcein for 1 h at 4°C for 15 min. Dye transfer of both R18 and calcein to large syncytia was seen in cells coexpressing most of the mutant F proteins with SER virus HN and incubated at 37°C (Fig. 3). The extent of lipid mixing and the number of

![Figure 4: Calcein transfer of I310M, N529K, and L511A mutants at various temperatures.](image-url)
FIG. 5. Effect of mutations on cell fusion activity. (A) Cell fusion assays. Dishes (35-mm diameter) of subconfluent HeLa T4 cells were infected for 1 h with the vTF7-3 virus at an MOI of 10 and then cotransfected with 3 μg of a wt or mutant SER virus F construct and a wt SER virus HN plasmid by the use of Lipofectin. At 16 to 20 h posttransfection, the cells were monitored for syncytium formation under a phase-contrast microscope. (B) Reporter gene activation assay. One population of HeLa T4 cells was infected with a recombinant vaccinia virus and transfected with a wt SER virus HN and a series of F mutants. The other cell population was infected with the wt vaccinia virus and transfected with a reporter gene construct. At 16 h posttransfection at 31°C, the two cell populations were mixed at a density of 10⁶ cells/ml in a 96-well tissue culture plate and incubated for 5 h at 37°C, after which cell fusion was quantitated by a colorimetric lysate assay. The data are percentages of the β-Gal activity observed in wt SV5 F-HN-expressing cells. The results are averages of three independent experiments.
content-mixing events were found to be fairly coincident in most of the mutants studied. The M370V mutant showed very low levels of R18 dye transfer and little or no calcein transfer at any of the temperatures studied. These results indicate that the N529K and I310M ectodomain mutants can substantially overcome the suppressive effect of the long SER virus F protein CT, whereas other ectodomain mutants, such as T149M, M489I, and T516V, also enhance fusion by the SER virus F protein, but to a lesser extent.

**Effect of N529K mutation on another SER virus mutant F protein and on mutant chimeric SV5/SER virus F proteins.** Since the effect of the CT mutation N529K on lipid mixing and content mixing was the most dramatic, we further studied the effect of this mutation in the context of a previously described SER virus F mutant, L539,548A, as well as a chimeric SV5/SER virus F protein carrying the CT from SER virus F or a chimeric SV5/SER-L539,548A,N529K mutant (33). All of the N529K mutants were expressed well and were transported to the cell surface at levels comparable to those of the respective original proteins (Fig. 6). The N529K mutants were found to be less stable with a 120-min chase period than the original mutant proteins.

**DISCUSSION**

SER virus, a paramyxovirus, exhibits minimal fusion activity, and previous studies demonstrated that an extended CT domain in its F protein plays an important role in fusion suppression (34). We also previously demonstrated that various site-specific mutations in the CT of the SER virus F protein can the β-Gal activity of wt SV5 F-HN. As shown in Fig. 5B, mutants K132E and P443S showed β-Gal activities ranging from 2 to 16% that of the wt SV5 F protein, whereas the T149A, I310M, T438S, M489I, and T516V mutants showed β-Gal activities of 25 to 54% that of the wt. The M370V mutant showed no β-Gal activity. The N529K mutant showed a fusion activity of 116% compared to the wt SV5 F protein coexpressed with the SV5HN protein. The L511A mutant served as a negative control, as it showed no β-Gal activity. These results indicate that the N529K and I310M ectodomain mutants can substantially overcome the suppressive effect of the long SER virus F protein CT, whereas other ectodomain mutants, such as T149M, M489I, and T516V, also enhance fusion by the SER virus F protein, but to a lesser extent.

**Cell fusion activity of mutant F proteins.** To investigate the effects of the mutations on the induction of syncytium formation, we coexpressed the mutant F proteins with the SER virus HN protein. As shown in Fig. 5, most of the ectodomain mutants induced extensive syncytium formation. The extent of syncytium formation was quantitated by a reporter gene activation assay (23), and the results were expressed as the percentages of β-Gal activity of the mutant F proteins relative to that of the wt F-HN. As shown in Fig. 5B, most of the ectodomain mutants showed β-Gal activities ranging from 2 to 16% that of the wt SV5 F protein, whereas the T149A, I310M, T438S, M489I, and T516V mutants showed β-Gal activities of 25 to 54% that of the wt. The M370V mutant showed no β-Gal activity. The N529K mutant showed a fusion activity of 116% compared to the wt SV5 F protein coexpressed with the SV5 HN protein. The L511A mutant served as a negative control, as it showed no β-Gal activity. These results indicate that the N529K and I310M ectodomain mutants can substantially overcome the suppressive effect of the long SER virus F protein CT, whereas other ectodomain mutants, such as T149M, M489I, and T516V, also enhance fusion by the SER virus F protein, but to a lesser extent.

**Effect of N529K mutation on another SER virus mutant F protein and on mutant chimeric SV5/SER virus F proteins.** Since the effect of the CT mutation N529K on lipid mixing and content mixing was the most dramatic, we further studied the effect of this mutation in the context of a previously described SER virus F mutant, L539,548A, as well as a chimeric SV5/SER virus F protein carrying the CT from SER virus F or a chimeric SV5/SER-L539,548A,N529K mutant (33). All of the N529K mutants were expressed well and were transported to the cell surface at levels comparable to those of the respective original proteins (Fig. 6). The N529K mutants were found to be less stable with a 120-min chase period than the original mutant proteins. As shown in Fig. 7, both lipid mixing and calcein transfer were enhanced to a large extent in the L539,548A,N529K and chimeric SV5/SER-L539,548A,N529K mutants; however, the fusion activity of a chimeric SV5/SER virus F mutant harboring the N529K mutation was enhanced to a lesser extent. The effect of the N529K mutation on lipid mixing and content mixing was also studied at low temperatures, and it was found that the L539,548A,N529K and chimeric SV5/SER-L539,548A,N529K mutants were able to induce dye transfer at temperatures as low as 18°C, whereas neither the original SER virus F mutant L539,548A nor the chimeric SV5/SER virus F protein showed dye transfer at reduced temperatures. The L539,548A,N529K, chimeric SV5/SER-N529K, and chimeric SV5/SER-L539,548A, N529K mutants also showed enhanced syncytium formation compared to the original mutant proteins. The chimeric SV5 F mutant L539,548A,N529K showed syncytia at 10 to 12 h posttransfection and was found to be highly fusogenic, with large multinucleate cells seen in all fields. In comparison, the SER virus F mutant L539,548A,N529K showed no syncytium formation until 16 h posttransfection (Fig. 8). The introduction of the N529K mutation also resulted in an increase in fusion, as assessed by a reporter gene activation assay (data not shown). This chimeric SV5/SER-L539,548A,N529K mutant showed the highest fusion activity of any of the constructs studied, suggesting that Asn 529 is one of the key residues involved in modulating fusion by the SER virus CT.

**DISCUSSION**

SER virus, a paramyxovirus, exhibits minimal fusion activity, and previous studies demonstrated that an extended CT domain in its F protein plays an important role in fusion suppression (34). We also previously demonstrated that various site-specific mutations in the CT of the SER virus F protein can...
overcome the fusion-suppressive effect of the long CT domain (33, 34). SER virus is very closely related to SV5 serologically and in its protein profile; however, besides the presence of the additional 22 amino acids in the CT, there are 9 amino acid differences spanning the ectodomain, the TM domain, and the CT domain between the SER virus and SV5 F proteins. Since the long CT of the SER virus F protein could only partially suppress fusion in a chimeric SV5 F protein carrying the CT of the SER virus F protein (33), we further studied the possible effects of these amino acid differences on fusion activity by mutating individual SER virus F residues to the corresponding SV5 residues.

We observed that the ectodomain mutants showed lipid mixing, calcein transfer, and syncytium formation to various extents. We performed our assays at a physiological pH to determine if these mutations could induce fusion under conditions in which the wt SER virus F protein shows no fusion. The SER virus F protein, when expressed without HN, does not fuse. However, with the addition of HN, it induces fusion. Since the long CT of the SER virus F protein is not sufficient to fully suppress fusion, we hypothesized that by mutating the CT of the SER virus F protein, we could enhance its ability to suppress fusion. Therefore, we studied the possible effects of introducing changes in residues in the CT of the SER virus F protein.
not show fusion at low pHs (32). Many of the mutant proteins also showed dye transfer at a less-than-optimal temperature for fusion (32, 25, and 18°C), indicating that these mutations lower the energy requirement for the SER virus F protein to induce fusion. It was shown previously that the presence of prolines at residues 22 and 443 destabilizes the SV5 F protein and decreases the energy requirement for triggering the conformational change to the fusion active state (25, 35). In contrast, in the wt SER virus F protein, prolines are present at both position 22 and position 443, but the protein is nonfusogenic. The P443S mutant showed some lipid mixing and calcein transfer at 37 and 32°C, although limited syncytium formation was observed at these temperatures. Taken together, these results indicate that the effect of the proline-to-serine change at residue 443 depends on the sequence context of the F protein.

We observed extensive dye transfer and syncytium formation when the Asn residue at position 529 was mutated to lysine, and fusion was found to be very dramatic, with extensive multinucleate cells. It is interesting that this single residue, also in the CT domain, can overcome the fusion-suppressive effect of the long CT. We also found that this mutation further enhanced the fusion activity of fusogenic mutants that we described earlier, such as the SER virus F L539,548A and chimeric SV5/SER virus F or chimeric SV5/SER-L539,548A F proteins (33). Surprisingly, the fusion observed with the SER-N529K mutant was more profound than that with the chimeric SV5/SER-N529K mutant. This indicates an effect of differences in the overall context of the SER virus F ectodomain compared with that of SV5. A highly fusogenic state was observed with the chimeric SV5/SER-L539,548A,N529K mutant, which had enhanced fusion kinetics (with syncytium formation observed as early as 12 h posttransfection) and showed fusion to a greater extent, suggesting a lower activation energy re-

FIG. 8. Effect of N529K mutation on fusion activity. Dishes (35-mm diameter) of subconfluent HeLa T4 cells were infected for 1 h with vTF7-3 at an MOI of 10 and then cotransfected with the mutant chimeric SV5/SER-L539,548A,N529K and SER-L539,548A,N529K F constructs and a wt SV5 HN or SER virus HN plasmid by the use of Lipofectin. The cells were monitored for syncytium formation under a phase-contrast microscope at different times posttransfection. Magnification, ×200.
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sion in the absence of the SER virus HN protein, in contrast to the
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the external domain of the SER virus F protein
the fusion glycoprotein of Newcastle disease virus suggests a novel paradigm
the fusion glycoprotein of pneumonia virus of mice suggests
the fusion glycoprotein of Sendai virus: interaction with neuraminic acid in affinity
the fusion glycoprotein of Sendai virus fusion protein (f) involves a conformational change with exposure of a new
the fusion glycoprotein of Sendai virus fusion protein inhibits virus-cell fusion. A plausible mode of


